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(57) Abstract

A bacterial cell (preferably a gram-negative, enteric bacterium such as V. cholerae) the chromosome of which contains a DNA sequence encoding a heterologous antigen, which sequence is functionally linked to an iron-regulated promoter such as the *irgA* promoter of V. cholerae.

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HETEROLOGOUS ANTIGENS IN LIVE CELL VACCINE STRAINS

The field of the invention is genetically engineered live bacterial cell vaccine strains.

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Statement as to Federally Sponsored Research

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Background of the Invention

V. cholerae is a gram-negative bacterium that causes a severe, dehydrating and occasionally fatal diarrhea in humans. There are an estimated 5.5 million cases of cholera each year, resulting in greater than 100,000 deaths (Bull. W.H.O. 68:303-312, 1990). Over the last several decades, cholera has been considered to occur primarily in developing countries of Asia and Africa, but recently it has reached epidemic proportions in regions of South and Central America, as well (Tauxe et al., J. Am. Med. Assn. 267:1388-1390, 1992; Swerdlow et al., J. Am. Med. Assn. 267:1495-1499, 1992).

Patients who recover from cholera infection have long-lasting, perhaps lifelong, immunity to reinfection (Levine et al., J. Infect. Dis. 143:818-820, 1981). The development of V. cholerae vaccines has focused on reproducing this naturally occurring immunity, but the currently available parenteral, killed whole-cell vaccine preparation provides less than 50% protection from disease, for a duration of only 3 to 6 months (Saroso et al., Bull. W.H.O. 56:619-627, 1978; Levine et al., Microbiol. Rev. 47:510-550, 1983). A genetically-engineered, live oral vaccine for V. cholerae has several theoretical advantages over the present parenteral killed whole-cell vaccine. As a mucosal pathogen, V. cholerae

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adheres selectively to the M cells of the gastrointestinal tract (Owen et al., J. Infect. Dis. 153:1108-1118, 1986) and is a strong stimulus to the common mucosal immune system (Svennerholm et al., Lancet 5 i:305-308, 1982); and oral cholera vaccination in humans produces a strong salivary gland IgA response to cholera toxin B subunit (Czerkinsky et al., Infect. Immun. 59:996-1001, 1991). Oral vaccines take advantage of the fact that oral administration of antigens appears to be 10 the most efficient stimulus for the development of secretory IgA (Svennerholm, supra), and that secretory IgA by itself is sufficient to protect against intestinal disease from V. cholerae (Winner III, et al., Infect. Immun. 59:977-982, 1991). Oral, killed whole cell 15 vaccines with or without the B subunit of cholera toxin have undergone extensive testing in volunteer and field trials over the past decade, and have been found to be more immunogenic and confer longer protection than the parenteral killed whole-cell vaccine (Svennerholm et al., 20 J. Infect. Dis. 149:884-893, 1984; Black et al., Infect. Immun. 55:1116-1120, 1987; Clemens et al., Lancet i:1375-1378, 1988; Clemens et al., J. Infect. Dis. 158:60-69, 1988; Jertborn et al., J. Infect. Dis. 157:374-377, 1988; Sack et al., 164:407-11, 1991).

Such killed whole-cell vaccines were traditionally favored over live whole-cell vaccines because the latter, which can multiply in the gut of the vaccinated animal, were considered unsafe. However, unlike killed-cell vaccines, live-cell vaccines would not require multiple doses, and in a rabbit model, live bacteria are more effective immunogens for secretory IgA than dead organisms (Keren et al., J. Immunol. 128:475-479, 1982). Live vaccines have the further advantage of potentially being transmitted from recipients to others in the

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The most important virulence factor for V. cholerae in causing clinical disease is cholera toxin, a protein complex consisting of one A subunit and 5 B subunits. Live, oral vaccine strains currently being 5 tested bear mutations in either the A subunit or in both subunits of cholera toxin (Mekalanos et al., Nature 306:551-557, 1983; Herrington et al., J. Exp. Med. 168:1487-1492, 1988; Levine et al., Lancet ii:467-470, 1988). An internal deletion of the gene encoding the A 10 subunit of cholera toxin (ctxA) in the classical strain 0395 produces a strain (0395-N1) which is highly immunogenic in humans, but produces non-specific symptoms in about half of the recipients (Mekalanos, supra; Herrington, supra; Mekalanos, U.S. Patent No. 4,882,278, 15 herein incorporated by reference), an indication that the strain is still virulent.

Summary of the Invention

As described in detail below, it has now been found that a V. cholerae gene, such as the irgA locus of 20 V. cholerae, can function as a site for the integration and high-level expression of sequences encoding heterologous antigens in vaccine strains of V. cholerae. IrgA, the major iron-regulated outer membrane protein of V. cholerae, is a virulence factor for this organism that 25 is independent of cholera toxin (Goldberg et al., USSN 07/629,102, herein incorporated by reference; Goldberg et al., Infect. Immun. 58:55-60, 1990). In vivo-grown V. cholerae expresses iron-regulated proteins that are not seen following growth in normal in vitro 30 conditions (Sciortino et al., 42:990-996, 1983), suggesting that the organisms sense low-iron conditions in the intestine. A mutation in irgA produces a 100-fold defect in the virulence of V. cholerae in a suckling mouse model. Regulation of irgA expression by iron is

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exceptionally tight, with a 1000-fold induction ratio in low- compared with high-iron conditions (Goldberg et al., Infect. Immun. 58:55-60, 1990). The entire structural gene of irgA has been cloned from the classical V. 5 cholerae strain 0395 (Goldberg et al., Mol. Microbiol. 6:2407-2418, 1992). Use of such an iron-regulated promoter to control expression of a heterologous antigen in a live vaccine strain has a number of distinct advantages. A high induction ratio ensures that the gene 10 encoding the heterologous antigen (1) will be expressed in the low-iron environment of the vaccinee's gut at a level high enough to ensure that it induces an immune response, and yet (2) will be expressed minimally when the cells are cultured in vitro, where high-level 15 expression would potentially provide selection pressure favoring inactivation of the gene and complicate largescale culturing of the cells necessary for vaccine production. Where, as in the case of irgA, the protein encoded by the naturally-occurring gene is, for at least 20 some V. cholerae strains, a virulence factor that is not essential for growth of the bacterium, insertion of the heterologous antigen coding sequence next to the promoter can be readily accomplished in such a way as to delete or otherwise inactivate the virulence factor coding 25 sequence, thereby decreasing the virulence of the engineered strain without affecting its viability.

The invention thus includes a genetically engineered V. cholerae chromosome containing a DNA sequence encoding a heterologous antigen, the DNA sequence being functionally linked to a naturally-occurring V. cholerae promoter. The heterologous antigen, defined as a polypeptide which is not expressed by the wildtype host species, is preferably a nontoxic polypeptide which is part or all of a protein that is naturally expressed by an infectious organism, and which

induces an antigenic response in an animal (preferably a mammal such as a human, non-human primate, cow, horse, sheep, goat, pig, dog, cat, rabbit, rat, mouse, guinea pig, or hamster). The infectious organism from which the 5 heterologous antigen is derived may be, for example, a bacterium, a virus, or a eukaryotic parasite, and the heterologous antigen may be, e.g., an OSP (Outer Surface Protein) of Borelia burgdorferai; animmunogenic, nontoxic subunit or fragment of a bacterial toxin such as Shiga 10 toxin, diphtheria toxin, Pseudomonas exotoxin A, pertussis toxin, tetanus toxin, anthrax toxin, one of the E. coli heat-labile toxins (LTs), one of the E. coli heat-stable toxins (STs), or one of the E. coli Shigalike toxins; an immunogenic portion of a viral capsid 15 from a virus such as human immunodeficiency virus (HIV), any of the Herpes viruses (e.g., Herpes simplex virus or Epstein-Barr virus), influenza virus, poliomyelitis virus, measles virus, mumps virus, or rubella virus; or an immunogenic polypeptide derived from a eukaryotic 20 parasite, such as the causative agent for malaria, pneumocystis pneumonia, or toxoplasmosis. (One preferred example of such a polypeptide is a malarial circumsporozoite protein.) By "functionally linked to a naturally-occurring V. cholerae promoter" is meant that 25 expression of the sequence encoding the heterologous antigen is controlled by a promoter which is found in wild-type V. cholerae, such as the ctxA promoter, or an iron-regulated promoter such as that of irgA. Construction of such a functional linkage can be 30 accomplished as described in detail below, or generally, using standard methods, by locating the desired promoter sequence sufficiently near to (and typically, though not necessarily, just upstream of) the promoterless heterologous antigen-encoding sequence to permit the 35 desired promoter sequence to control expression of the

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latter sequence. Functional siting of promoter sequences is well within the abilities of one of ordinary skill in the art of prokaryotic gene expression. Where the promoter naturally controls the expression of a V. 5 cholerae virulence factor that is nonessential for growth of the cell, the sequence encoding that virulence factor will preferably be deleted or otherwise mutated to prevent expression of a biologically active form of that virulence factor. Preferably, the ctxA locus on the 10 chromosome will also be deleted or otherwise inactivated, so that biologically active cholera toxin cannot be expressed from the chromosome. Such deletions, mutations and insertions can readily be carried out by one of ordinary skill using the methods described herein, or 15 other well-known, standard techniques. In preferred embodiments, the ctxA deletion is identical to that of strain 0395-N1 (Mekalanos, U.S. Patent No. 4,882,278).

Also within the invention is a bacterial chromosome (preferably from a gram-negative, enteric 20 bacterium such as V. cholerae), containing a DNA sequence encoding a heterologous antigen, which sequence is functionally linked to an iron-regulated promoter which functions in the host bacterium to permit significantly (i.e., at least ten-fold and preferably 100-fold) higher 25 expression of the heterologous antigen in a low-iron environment, such as in an animal's intestine, than in a high-iron environment, such as under typical in vitro culture conditions. An example of such a promoter is the naturally-occurring promoter of V. cholerae irgA, which 30 includes at a minimum a sequence substantially identical to nucleotides 1000 through 1041 (SEQ ID NO: 2), inclusive, of the sequence shown in Fig. 5 (SEQ ID The promoter sequence used is preferably nucleotides 922 to 1041 (SEQ ID NO: 3), more preferably 35 922 to 1079 (SEQ ID NO: 4) or 1000 to 1079 (SEQ ID

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NO: 5), still more preferably 905 to 1041 (SEQ ID NO: 6) or 905 to 1079 (SEQ ID NO: 7), and most preferably 905 to 1438 (SEQ ID NO: 8), 922 to 1438 (SEQ ID NO: 9), or 1000 to 1438 (SEQ ID NO: 10) (all inclusive). Examples of 5 other iron-regulated promoters which would be useful in the invention are those derived from the fatA gene of V. anquillarum (Koster et al. J. Biol. Chem. 266:23829-23833, 1991); E. coli slt-IA (or other E. coli Furbinding promoter sequences, as discussed by Calderwood et al., J. Bacteriol. 169:4759-4764, 1987; De Grandis et al., J. Bacteriol. 169:4313-4319, 1987; and DeLorenzo et al., J. Bacteriol. 169:2624-2630, 1987); the iron-regulated outer membrane proteins of Salmonella typhi (Fernandez et al., Infect. Immun. 57:1271-1275, 1989), 15 the iron-regulated hemolysin promoter of Serratia (Poole

- the iron-regulated hemolysin promoter of Serratia (Poole et al., Infect. Immun. 56:2967-2971, 1988); the Yersenia iron-regulated promoters (Carniel et al., Molecular Microbiol. 6:379-388, 1992; Staggs et al., J. Bacteriol. 173:417-425, 1991; and Staggs et al., Molecular
- 20 Microbiol. 6:2507-2516, 1992); the V. vulnificus ironregulated promoters; the Pseudomonas exotoxin A ironregulated promoter (Bjorn et al., Infect. Immun. 19:785791, 1978); and Plesiomonas iron-regulated genes involved
 in heme-iron uptake (Daskaleros et al., Infect. Immun.
- 25 59:2706-2711, 1991). It is believed that most if not all enteric, gram-negative bacterial species, including E. coli, Salmonella, Shigella, Yersenia, Citrobacter, Enterobacter, Klebsiella, Morganella, Proteus, Providencia, Serratia, Vibrios, Plesiomonas, and
- 30 Aeromonas, utilize highly similar fur-binding, ironregulated promoter sequences, and it is likely that they
 also utilize secondary iron-regulated promoter sequences
 similar to that of *irgA*. Such promoter sequences are
 well-known to those of ordinary skill, or can be readily
- 35 determined from current information regarding iron-

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regulated promoters. Construction of such promoter sequences adjacent to a given heterologous antigenencoding sequence, and insertion of the resulting construct into a V. cholerae genome, is readily accomplished by one of ordinary skill; the ability of such a promoter to function as predicted can then be tested in low- and high-iron conditions as described below, without undue experimentation.

Also within the invention is a V. cholerae cell, 10 or a homogeneous population of such cells, which contains the genetically engineered chromosome described above. Such cells can be said to define a vaccine strain useful, when combined with a pharmaceutically acceptable diluent suitable for oral administration, as a live-cell vaccine. 15 Administration of such a vaccine to an animal (e.g., a human or other mammal) will provoke immunity not only to V. cholerae, but also to an antigen derived from a second organism; it thus serves as a bivalent vaccine. An example of such a vaccine utilizes a genetically 20 engineered V. cholerae strain in which the ctxA and irgA coding sequences are largely deleted and a sequence encoding Shiga-like toxin B subunit is functionally linked to the irgA promoter. This strain is described in detail below. Of course, the bacterial strain of the 25 invention could be engineered to encode several heterologous antigens, each linked to an identical or different iron-regulated promoter, to produce a multivalent vaccine effective for simultaneously inducing immunity against a number of infectious diseases.

Other features and advantages will be apparent from the detailed description provided below, and from the claims.

Brief Description of the Drawings

Fig. 1 is a schematic diagram illustrating the construction of plasmids used in this study. A partial restriction map of 0395 chromosomal DNA is shown with 5 relevant restriction enzyme sites, using base-pair numbering as in Goldberg et al., Mol. Microbiol. 6:2407-2418, 1992; and Goldberg et al., Proc. Natl. Acad. Sci. USA 88:1125-1129, 1991. The location of irgA, the location of fragments cloned in the construction of vaccine strains and the locations of fragments used as probes in Southern blot analysis are indicated. The upstream irgA fragment is indicated by a solid bar; the downstream irgA fragment by a hatched bar; and the slt-IB subunit fragment by a stippled bar. Plasmids and chromosomal fragments are not drawn to scale.

Figs. 2A-2B is a set of Southern blots
illustrating hybridization of chromosomal DNA from wildtype and mutant V. cholerae strains, digested with
HindIII, separated by agarose electrophoresis and probed
20 as follows: (A) SmaI - HincII fragment (region deleted
in vaccine strains); (B) HincII - HincII fragment
(downstream probe); (C) HindIII - SmaI fragment (upstream
probe); (D) EcoRV - HindIII fragment from pSBC52 (slt-IB
subunit probe). Lanes: 1, 0395-N1; 2, SBC20; 3, B014-1;
25 4, B024-1; 5, VAC1; 6, VAC2; 7, 0395-N1. The genomic
location of the fragments used as probes is indicated in
Fig. 1. The numbers to the left of the blot indicate the
sizes (in kbp) of DNA standards.

Fig. 3 is a photograph of an SDS-PAGE analysis of the outer membrane proteins expressed by certain V. cholerae strains when grown in high- or low-iron medium. Lanes: 1, 0395-N1 grown in high-iron medium; 2, 0395-N1 grown in low-iron medium; 3, SBC20 grown in low-iron medium; 4, VAC1 grown in low-iron medium; 5, VAC2 grown in low-iron medium; 6, 0395-N1 grown in low-iron medium.

The numbers to the left of the gel indicate the molecular masses (in kDa) of the protein standards.

Fig. 4 is a schematic diagram of the construction of the pSBC52 plasmid utilized in these experiments.

- 5 pSBC32 (Calderwood et al., Infect. Immun. 58:2977-2982, 1990) was subjected to PCR using primer No. 1: 5'-CCGAATTCTCTAGAGATATCGTGTGGAATTGTGAGCGGATAA-3' (SEQ ID NO: 11), which introduces restriction sites for EcoRI, XbaI, and EcoRV, and primer No. 2:
- 10 5'-CCAAGCTTCTGCAGCCCGGGATTTAACATTTATGAATCTCCGCCT-3' (SEQ ID NO: 12), which introduces restriction sites for HindIII, PstI, and SmaI. The PCR product was then digested with EcoRI and HindIII, and cloned into EcoRI/HindIII-digested pUC19, to produce pSBC52.
- 15 Fig. 5 shows the nucleotide sequence of a portion of the *irgA* cDNA (SEQ ID NO: 1), including the promoter sequence. A 19-bp interrupted dyad symmetric element homologous to the Fur box of *E. coli* is indicated by inverted horizontal arrows below the sequence. Vertical 20 lines mark the margins of what is believed to be regions

important for irgA promoter function.

<u>Detailed Description</u>

In the experiments described below, the non-toxic B subunit of Shiga toxin was used as a model heterologous antigen, because of the easily available assays for this protein (Donohue-Rolfe et al., J. Clin. Microbiol. 24:65-68, 1986), as well as the possible role that antibodies against the B subunit play in protecting against severe Shigellosis and hemolytic uremic syndrome. Shiga toxin is a heterodimeric protein consisting of one A subunit (MW 32 kDa) and five B subunits (MW 7.7 kDa) (Seidah et al., J. Biol. Chem. 261:13928-13931, 1986); the B subunit of Shiga toxin is identical in amino acid sequence to the B subunit of Shiga-like toxin I produced by

enterohemorrhagic strains of E. coli (Calderwood et al., Proc. Natl. Acad. Sci. USA 84:4364-4368, 1987). This identical protein product is referred to as StxB throughout this study. Immune response to Shiga toxin is primarily directed against the B subunit, and antibodies directed against this subunit, or against synthetic peptides from regions of the subunit, provide protective immunity against holotoxin (Donohue-Rolfe et al., J. Exp. Med. 160:1767-1781, 1984; Harari et al. Infect. Immun. 56:1618-1624, 1988; Harari et al., Mol. Immunol. 27:613-621, 1990; Boyd et al., Infect. Immun. 59:750-757, 1991).

Described below are the insertion of a promoterless gene for the Shiga-like toxin I B subunit (slt-IB) into an irgA deletion, and the introduction of this construct into the chromosome of the V. cholerae ctxA deletion strain 0395-N1, thus producing a live, attenuated vaccine strain of V. cholerae that contains StxB under the transcriptional control of the iron-regulated irgA promoter.

20 MATERIALS AND METHODS Bacterial strains and plasmids.

The bacterial strains and plasmids used in this study are described in Table 1, with the exception of plasmids pMBG126, pSAB18, pSAB12, pSAB19, pSAB14, and pSAB24, which are described in detail below and are depicted in Fig. 1; and plasmid pSBC52, which is described in the description of Fig. 4 provided above. Standard plasmid cloning vectors pUC18, pUC19, and pBR322 are commercially available (e.g., Pharmacia).

30 Media.

All strains were maintained at -70°C in Luria broth (LB) media (Sambrook et al., A Laboratory Manual, 2nd ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989), containing 15% glycerol. LB media,

with or without the addition of the iron chelator 2,2-dipyridyl (final concentration, 0.2 mM), was used for growth in low- and high-iron conditions, respectively. Ampicillin (100 μ g/ml), kanamycin (45 μ g/ml), and streptomycin (100 μ g/ml) were added as appropriate. Genetic methods.

DNA, restriction enzyme digests, agarose gel electrophoresis, and Southern hybridization of DNA

10 separated by electrophoresis were performed according to standard molecular biologic techniques (Sambrook, supra). GeneScreen Plus hybridization transfer membranes (DuPont Biotechnology Systems, NEN Research Products, Boston, MA) were used according to the manufacturer's protocols for Southern hybridization. DNA sequencing was performed using the Sequenase DNA Sequencing Kit (United States Biochemical Corporation, Cleveland, OH).

Plasmids were transformed into *E. coli* strains by standard techniques, or were electroplated into *V*.

20 cholerae using a Gene Pulser (Bio-Rad Laboratories, Richmond, CA), following the manufacturer's protocol, and modified for electroporation into *V. cholerae* as previously described (Goldberg et al., Proc. Natl. Acad. Sci. USA 88:1125-1129, 1991). Electroporation conditions were 2,500 V at 25-μF capacitance, producing time constants of 4.7-4.9 ms.

DNA restriction endonucleases, T_4 DNA ligase, calf intestinal alkaline phosphatase, and the Klenow fragment of DNA polymerase I were used according to the 30 manufacturers' specifications. Restriction enzymedigested chromosomal and plasmid DNA fragments were separated on 1% agarose gels; required fragments were cut from the gel under ultraviolet illumination and purified by electroelution (Sambrook et al, 1989, supra). DNA fragments used as probes were radiolabeled with α -32p-dCTP

using a random priming labeling kit (Prime Time "C" Oligonucleotide Labeling Biosystem, International Biotechnologies, Inc., New Haven, CT).

Construction of plasmids.

DNA was recovered upstream and at the 5' terminus of irgA as a HindIII-SmaI fragment from pMBG59, which contains the irgA promoter (irgP) (Goldberg et al., J. Bacteriol. 172:6863-6870, 1990) (Fig. 1). This fragment was cloned into the HindIII and SphI sites of pUC18 to 10 yield plasmid pMBG126; the SphI site of pUC18 had first been made blunt-ended by treatment with mung bean nuclease. DNA sequence analysis of pMBG126 revealed that the SphI site was unexpectedly preserved at the junction with SmaI; the sequence was otherwise as predicted. 15 was then recovered at the 3' terminus and downstream of irgA as a 1.5 kilobase-pair (kbp) HincII fragment from plasmid pSAB25. SacI linkers were added to this fragment and it was ligated into the unique SacI site of pMBG126, in the same orientation as the upstream irgA fragment, to 20 yield plasmid pSAB18. The internal SalI site in the pUC polylinker of pSAB18 was removed by digesting with SalI, treating with the Klenow fragment of DNA polymerase I, and religating the blunt ends, to create pSAB12. A DNA segment encoding the promoterless B subunit of Shiga-like 25 toxin I (slt-IB) was recovered as an EcoRV-SmaI fragment from plasmid pSBC52. This fragment was introduced into the unique EcoRV and Smal sites of pSAB18, such that slt-IB was under the transcriptional control of irgP on the upstream irgA fragment, yielding plasmid pSAB19. 30 construction of plasmids pMBG126, pSAB18, pSAB12, and pSAB19 was verified by restriction enzyme digestion and double-stranded DNA sequencing.

The desired fragments were then introduced into the suicide vector pCVD442 as follows. pSAB12 and pSAB19

35 were digested with HindIII and EcoRI and the DNA fragment

containing either the *irgA* deletion (from pSAB12) or the *irgA* deletion-slt-IB-substitution (from pSAB19) were made blunt-ended by the Klenow fragment of DNA polymerase I. Following ligation to SalI linkers, the fragments were ligated into the unique SalI site of pCVD442, yielding plasmids pSAB14 and pSAB24 respectively, and propagated in the permissive strain SM10 & pir. Plasmid pCVD442 is a recently described suicide vector containing the pirdependent R6K replicon, ampicillin resistance, and the sacB gene from Bacillus subtilis (Donnenberg et al., Infect. Immun. 59:4310-4317, 1991).

Construction of VAC1 and VAC2

V. cholerae strain SBC20 is an irgA::TnphoA derivative of 0395-N1 (Pearson et al., Res. Microbiol. 15 141:893-899, 1990). The kanamycin resistance marker in TnphoA allowed screening of mutants for deletion of irgA (and hence TnphoA) by assessing susceptibility to kanamycin. The irgA allele of SBC20 was replaced with either the previously constructed irgA deletion, or the 20 irgA deletion containing slt-IB, as follows. Plasmids pSAB14 and pSAB24 were electroporated into SBC20, with selection for ampicillin and streptomycin resistance. Doubly-resistant colonies contained the respective plasmids integrated into the chromosome by homologous 25 recombination involving either the upstream or downstream fragments of irgA on pSAB14 or pSAB24, with creation of a merodiploid state. One such colony from the integration of pSAB14 into the chromosome of SBC20 was selected and named B014-1; one from the integration of pSAB24 into the 30 chromosome of SBC20 was named BO24-1. BO14-1 and BO24-1 were grown overnight in LB media without ampicillin selection, then plated on LB media with 10% sucrose but without NaCl, and grown at 30°C for 30 hours, thereby selecting for clones that had deleted the integrated sacB

35 gene (Blomfield et al., Mol. Microbiol. 5: 1447-1457,

1991). Sucrose-resistant colonies that are ampicillin susceptible but kanamycin resistant have re-excised the plasmid (yielding the parent SBC20, which contains the kanamycin resistance marker in TnphoA); those that are 5 both ampicillin and kanamycin susceptible have resolved the merodiploid state to replace the irgA locus in SBC20 with either the irgA deletion from pSAB14 or the irgA deletion-slt-IB fragment from pSAB24. Approximately 10% of sucrose-resistant colonies that were ampicillin-10 susceptible were also kanamycin-susceptible. One of these colonies which had replaced the irgA:: TnphoA locus with the irgA deletion was further purified and named VAC1; one which had replaced the irgA:: TnphoA locus with irgA::irgP-slt-IB was named VAC2. Confirmation of the 15 proper constructions in VAC1 and VAC2 was obtained by Southern hybridization of restriction enzyme-digested

20 irgA segment.

Preparation of outer-membrane proteins, whole cell
proteins, and periplasmic extracts.

chromosomal DNA that was probed with several different

DNA fragments to verify the expected deletion in irgA, as well as the introduction of the slt-IB within the deleted

Enriched outer membrane proteins were prepared from strains following growth in low- and high-iron media as previously described (Goldberg, Infect. Immun. 58:55-60, 1990). Proteins were separated by electrophoresis on a sodium dodecyl sulfate/10% polyacrylamide (SDS-PAGE) gel and visualized by staining with Coomassie brilliant blue. Whole cell proteins and periplasmic extracts were prepared from exponentially growing cells as previously described (Hovde et al., Proc. Natl. Acad. Sci. USA 85:2568-2572, 1988).

Immunodetection of StxB production.

Whole cell proteins and periplasmic extracts were 35 separated on a SDS-15% PAGE gel as described above, then

transferred to a NitroBind Transfer Membrane (Micron Separations Inc., Westboro, MA) with a semidry blotting apparatus (Hoefer Scientific Instruments, San Francisco, CA). Immunoreactive proteins were visualized after 5 sequential incubation with polyclonal rabbit anti-Shiga toxin antiserum and goat anti-rabbit IgG-conjugated alkaline phosphatase (Sigma Chemical Co., St. Louis, MO), followed by staining for phosphatase activity as described previously (Hovde, supra). The amount of StxB 10 present in periplasmic extracts or culture supernatants was quantitated with an enzyme-linked immunosorbent assay (ELISA) developed for the detection of Shiga toxin and modified for detection of purified StxB (Donohue-Rolfe et al., J. Clin. Microbiol. 24:65-68, 1986; Calderwood et 15 al., Infect. Immun. 58:2977-2982, 1990).

HeLa cell cytotoxicity.

The cytotoxicity of periplasmic extracts or culture supernatants for HeLa cells was assayed in a quantitative cytotoxicity assay by determining the extent 20 of HeLa cell detachment from microtiter plates (Gentry et al., J. Clin. Microbiol. 12:361-366, 1980). HeLa cells were grown at 37°C in a 5% CO₂ atmosphere in McCoy 5a (modified) medium containing 10% fetal calf serum and 100 μ g of penicillin and streptomycin per ml. Freshly 25 trypsinized cells were suspended in 0.1 ml of growth medium and allowed to attach to the wells of microtiter plates overnight. Serial dilutions of samples were added and the plates were again incubated overnight. The cells were fixed and then stained with crystal violet in a 5% 30 ethanol - 2% formaldehyde solution. Stained cell monolayers were dissolved in ethanol and the ${\tt A}_{{\tt 595}}$ read with a microtiter plate colorimeter.

Evaluation of virulence of vaccine strains.

50% lethal dose (LD_{50}) assays were performed by oral inoculation of 3- to 4-day old CD1 suckling mice

with either the parent V. cholerae strain 0395, an irgA mutant strain MBG40 (Goldberg et al., Infect. Immun. 58:55-60, 1990), the ctxA mutant strain 0395-N1, or VAC2. Cholera strains were grown overnight in LB medium at 30°C, pelleted, and resuspended in 0.5M NaHCO3 (pH 8.5). Mice were orally inoculated with serial dilutions of organisms, then kept at 30°C. Four or more mice were used per dose of bacteria. Survivial was determined at 40 h (Taylor et al., Proc. Natl. Acad. Sci. USA 84:2833-10 2837, 1987).

RESULTS

Confirmation of vaccine strain construction.

- (i) Southern hybridization analysis. To confirm the construction of the vaccine strains, chromosomal DNA 15 was purified from V. cholerae parent strains 0395-N1 and SBC20, the merodiploid strains B014-1 and B024-1, and the vaccine strains VAC1 and VAC2. The chromosomal DNAs were digested with HindIII, separated on agarose gels, and transferred to membranes for Southern hybridizations.
- 20 The Southern hybridizations of these digests, probed with four different fragment probes, are shown in Fig. 2. The location of the fragment probes within the *irgA* gene is shown in Fig. 1. The presence and size of the recognized fragments is consistent with the constructions depicted
- in Fig. 1, confirming the deletion of irgA in VAC1 and the deletion-replacement of the irgA locus with irgA::irgP-sltIB in VAC2.
- ii. Outer membrane protein analysis. Outer membrane proteins were prepared from strain 0395-N1 grown in low- and high-iron media and from strains SBC20, VAC1 and VAC2 following growth in low-iron media, then separated by electrophoresis on a SDS-PAGE gel (Fig. 3). IrgA, the 77 kilodalton (kDa) major iron-regulated outer membrane protein (Goldberg et al., Infect. Immun. 58:55-35 60, 1990), is present in 0395-N1 grown in low iron but is

absent in SBC20 (an *irgA* mutant) and the vaccine strains, confirming the deletion of *irgA* in VAC1 and VAC2.

Iron-regulated expression of the Shiga toxin B subunit in VAC2.

- (i) Western blot analysis of StxB production in VAC2. Western blot analysis of whole cell proteins and periplasmic extracts of VAC2 grown in high- and low-iron media demonstrated the production of a 7.7 kDa protein recognized by polyclonal rabbit anti-Shiga toxin
- antiserum in both whole cell proteins and periplasmic extracts prepared from VAC2 grown in low-iron media; no such protein was recognized in proteins prepared from the vaccine strain grown in high-iron media, demonstrating that the production of StxB is tightly iron-regulated
- 15 (data not shown).
 - (ii) <u>Quantitation of StxB production from irqP-slt-IB in plasmid pSAB19 and VAC2</u>. To verify iron-regulated production of StxB by *irqP-slt-IB* in plasmid pSAB19, and compare it with StxB production by VAC2, we
- first had to return pSAB19 to the V. cholerae background because irgP is not active in E. coli (Goldberg et al., Proc. Natl. Acad. Sci. USA 88:1125-1129, 1991). The production of StxB by strains 0395-N1(pSAB19) and VAC2 was quantitated using a sandwich ELISA, with a monoclonal
- 25 antibody specific for StxB as the capture molecule.
 Purified StxB, in measured amounts, was used as the
 standard. As shown in Table 2, both 0395-N1(pSAB19) and
 VAC2 express StxB in a tightly iron-regulated fashion, as
 expected, and produce five times the amount of B subunit
- 30 made by the reference strain, Shigella dysenteriae 60R, under low-iron conditions.

Virulence of vaccine strains.

(i) <u>Cytotoxicity to HeLa cells</u>. The cytotoxicity of periplasmic extracts or culture supernatants of
 35 strains 0395-N1(pSAB19) and VAC2, grown in low-iron

media, was assayed as described (Gentry et al., J. Clin. Microbiol. 12:361-366, 1980), and compared to the S. dysenteriae strain 60R. Neither 0395-N1(pSAB19) or VAC2 had detectable cytotoxicity in periplasmic extracts or supernatants, in contrast to periplasmic extracts of S. dysenteriae 60R, which were cytotoxic to at least a 10⁵-fold dilution (data not shown).

(ii) LD₅₀assays. The results of LD₅₀ assays for the wild-type V. cholerae strain 0395, ctxA mutant strain 0395-N1, irgA mutant strain MBG40, and vaccine strain VAC2 in the suckling mouse model are shown in Table 3. V. cholerae strain MBG40, an irgA::TnphoA mutant of strain 0395, had an LD₅₀ in suckling mice that was 2 orders of magnitude higher than that for the parental strain 0395, as previously demonstrated (Goldberg et al., Infect. Immun. 58:55-60, 1990). Strain 0395-N1, deleted for the A subunit of cholera toxin, was avirulent at an inoculum of 2 x 10⁹ organisms in this model. The vaccine strain VAC2, despite expressing StxB at high level,
20 remains avirulent in this model at an inoculum of 2 x 10⁹ organisms, similar to its parent strain 0395-N1.

USE

The *V. cholerae* strains of the invention are useful as bivalent vaccines capable of inducing immunity to *V. cholerae* and to an antigen derived from a second infectious organism. Because the strains are attenuated (i.e., do not induce a significant toxic reaction in the vaccinee), they can be used as live-cell vaccines, permitting effective immunity to result from administration of a single dose of the vaccine. An effective oral dose of the vaccine would contain approximately 10⁶ to 10⁸ bacteria in a volume of approximately 150 ml liquid. The diluent used would typically be water or an aqueous solution, such as

5

2 grams of sodium bicarbonate dissolved in 150 ml distilled water, which may be ingested by the vaccinee at one sitting, either all at once or over any convenient period of time.

Other Embodiments

Other embodiments are within the claims set forth below. For example, the host bacterium (the bacterium the chromosome of which is engineered to encode a heterologous antigen) can be E. coli or any other enteric 10 bacterium, including Salmonella, Shigella, Yersenia, Citrobacter, Enterobacter, Klebsiella, Morganella, Proteus, Providencia, Serratia, Plesiomonas, and Aeromonas, all of which are known or believed to have iron-regulated promoters similar to the Fur-binding 15 promoters of E. coli, and which may have other ironregulated promoters analogous to that of irgA. Also potentially useful would be a bacille Calmette-Guerin (BCG) vaccine strain engineered to encode a heterologous antigen linked to an iron-regulated promoter. 20 promoter used can be native to the species of the host bacterium, or can be a heterologous promoter (i.e., from a species other than that of the host bacterium) engineered into the host bacterium along with the heterologous antigen coding sequence, using standard 25 genetic engineering techniques. Multiple heterologous antigen coding sequences linked to the same or different iron-regulated promoter sequences can be inserted into a given chromosome, using techniques analogous to those set forth above, to produce a multivalent vaccine strain.

Those who practice in the field of prokaryotic gene expression will realize that, while naturally-occurring promoter sequences are preferred, synthetic sequences such as a consensus Fur-binding sequence or a hybrid of two or more Fur-binding sequences would also be

expected to be useful in the chromosomes of the invention. Alteration, addition or deletion of one or a few nucleotides within a naturally-occurring promoter sequence such as the *irgA* promoter would generally not affect its usefulness. The invention therefore encompasses iron regulated promoters having such inconsequential changes.

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Table 1. Bacterial strains and plasmids used in this study

Strain or plasmid	Relevant genotype or phenotype Ref. or	source
V. cholerae strain	8	
0395	Sm ^r	1
0395-N1	0395 ctxA, Sm ^r	1
SBC20	0395-N1 irgA::TnphoA, Smr, Kmr	2
MBG40	0395 irgA::TnphoA, Sm ^r , Km ^r	3
B014-1	SBC20 with pSAB14 integrated into	
	irgA, Sm ^r , Km ^r , Ap ^r	4
BO24-1	SBC20 with pSAB24 integrated into	
	irgA, Sm ^r , Km ^r , Ap ^r	4
VAC1	0395-N1 <i>\(\delta irg\text{A} \), \(\text{Sm}^r \)</i>	4
VAC2	0395-N1 AirgA::irgP-slt-IB, Sm ^r	4
E. coli strains		
SM10 λ pir	thi thr leu tonA lacY supE	5
	recA::RP4-2-Tc::Mu λ pirR6K, Km ^r	
Plasmids		
pMBG59	pBR322 with 4.7-kbp of V. cholerae	6
	MBG40 chromosome, containing DNA	
	upstream and at the 5' terminus	
	of irgA, as well as the	
	irgA:: TnphoA fusion joint from	
	this strain.	
pSAB25	3.0 kbp SmaI - MluI fragment of	4
	V. cholerae 0395 chromosome, containing	
	DNA at the 3'terminus and downstream	
	of irgA, made blunt-ended at the MluI	
	site and ligated into Smal-digested	
	pUC19.	
pSBC52	pUC19 with a promoterless gene for the	4
	B subunit of SLT-I (identical to StxB)	
	cloned between the EcoRI and HindIII	
	sites.	
pCVD442	Suicide vector composed of the mob,	7
-	ori, and bla regions from pGP704 and the	
	sacB gene of Bacillus subtilis.	

Apr, ampicillin resistance; Kmr, kanamycin resistance; Smr, streptomycin resistance.

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Ref. or source:

- 1. Mekalanos et al., Nature 306:551-557, 1983.
- 2. Pearson et al., Res. Microbiol. 141:893-899, 1990.
- 3. Goldberg et al., Infect. Immun. 58:55-60, 1990.
- 5 4. This study.
 - 5. Miller et al., J. Bacteriol. 170:2575-2583, 1988.
 - 6. Goldberg et al., J. acteriol. 172:6863-6870, 1990.
 - 7. Donnenberg and Kaper, Infect. Immun. 59:4310-4317, 1991.

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Table 2. Production of Shiga toxin B subunit by various strains following growth in high- and-low iron conditions

Strain	Periplasmi	.c extract ^a	Supernata	at ^a
5	High-iron	Low-iron	High-iron	Low-
iron				
0395-N1	b			
0395-N1 (pSAB19)	15.5	3,620	0.16	3.5
VAC2	0.87	4,130	45 45	0.73
S.dysenteriae 60R	238	674	0.8	16.4

a ng/50 OD₆₀₀ of original culture

b < 0.1 ng

- 25 **-**

Table 3. Virulence assays of wild-type and mutant strains of Vibrio cholerae in suckling mice

	Strain	LD ₅₀
5		(no. of bacteria)
	0395	1 x 10 ⁵
	MBG40	1 x 10 ⁷
	0395-N1	> 2 x 10 ⁹
0	VAC2	> 2 x 10 ⁹

- 26 -

SEQUENCE LISTING

(1)	GENERAL	INFORMATION:
-----	---------	--------------

(i) APPLICANT:

Calderwood, Stephen B.

Butterton, Joan R.

Mekalanos, John J.

(ii) TITLE OF INVENTION:

HETEROLOGOUS ANTIGENS IN LIVE CELL

VACCINE STRAINS

(iii) NUMBER OF SEQUENCES:

12

(iv) CORRESPONDENCE ADDRESS:

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U.S.A.

02110-2804

(v) COMPUTER READABLE FORM:

(A) MEDIUM_TYPE:

3.5" Diskette, 1.44 Mb

(B) COMPUTER:

IBM PS/2 Model 50Z or 55SX

(C) OPERATING SYSTEM:

MS-DOS (Version 5.0) WordPerfect (Version 5.1)

(D) SOFTWARE:

(vi) CURRENT APPLICATION DATA:

(A) APPLICATION NUMBER:

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(B) FILING DATE: (C) CLASSIFICATION: February 22, 1993

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- (A) APPLICATION NUMBER:
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(viii) ATTORNEY/AGENT INFORMATION:

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200154

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH:

(B) TYPE:

nucleic acid

(C) STRANDEDNESS:

both linear

(D) TOPOLOGY:

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

ATC	GATG	ATA	AAAA	ATCC	CG C	TGCG	CGG	G AT	TTTT	TATT	GCC	ACTC	ATC	GGGC	CTTGC	CT	60
TGG	CGGA	GCG	CATC	AATA	AA T	AGGC	GCAG	C CG	AAGT	GGT	GAC	GACC	GAG	CGGA	TAGAA	A G	120
CAG	TTGA!	TTT	CTGT	TGGC:	rg T	GATT	GCCA?	r cc	GTTG	ACGC	AAG	GAAT	GAG	GCTG	CCCGG	3A	180
TGC	CCG	TTT	CAAA	ACCA:	TT G	GCAA	ACCAI	A GT	GGGA	AGCA	AAC	CAAT	ACC .	ACGA	CCTT1	ra	240
GCA	ATCG	CAT	CGGC'	TTGC	AT G	GCAA	GATTA	A TC	GCTT	rgta	AAC	GACT	CTC	TAGT	GCTGG	3C	300
AGTO	GAAT	AAC	TGCC	GAAC:	rc T	GGAT	GTG	C AG	TTCA	AGCT	CCG	CGCG	CCG .	ACAA	GCAAI	ra	360
AAA	CAA!	rcc	ATGG	GTGA:	rg A	ATCA	GCTC	A CG	AGGA:	rggg	TCG	GTTT	ATC	TCGA'	TGGGC	cc	420
LAAA	TTTAT	rgg	GAGA	GCG:	TA A	GTGG	CATAC	G CG	CCAA!	TAGC	CTA	AGCG'	TTC '	TTTG	CGATA	LA	480
CCC	ATGG	GGG	CGGC	GTGT	rc A	atcci	AAATO	ATC	CAAA!	rcgg	GCT	CAAA	CAC	CTCA'	TCACT	rg ·	540
TGTI	[GAA]	ACT	GGCT	GAGT	AG A	CGGA!	CTTC	AA!	rgtc	GAAT	GCT	CTG	CAT	AAAC'	CATC	c	600
AATA	ACTTO	3GC	TGAG	CCAG	CC G	CGGA:	CAAI	A TTO	GGG:	rgta	CCA	CCAG	CGT	GAGT	rcccc	ZA.	660
GTC	CTT	GAT	TGTT	CAATI	rc T	TGCAI	ACGC	TC	CTGA	CTTT	TAT	rggc	CAG	TTCA	AGTAG	T	720
TGCI	rccgi	AGT .	AAAC	CGCA	AA C	ACTT	CTCCI	C GC	rttg	STGA	GCG	CTAAC	GCG (GTTG(CCTTG	A	780
CGC	TCA	ACA	AGCT:	TTGT	cc c	AAGT	CTC	TCI	AAGT:	rgcg	CCAI	AACG	GCG :	ACTC	AGGGI	G	840
GATI	TAGO	CT	GTTC!	AAGC	C T	TTGG	CAGCO	G GCZ	AGTC	AGGC	TCT	ratg:	rtg (GCAA	AGCGC	:A	900
TGG	AAG	CTT	TTAC	GCGC	CT G	AGATO	CTTGC	C AT	AGGT	TTT	GAC	CCTT	AAA	GAAT	ATTA	C	960
CACA	GAC	GTT	CCAT	ATTTC	GG A	CCGAI	ACTA1	TCC	CATG	CTC	ĠAT	CTATO	CTC (CAGT	ACAGA	LA	1020
TATA	TGA	ATA :	ATCC	CTT	CT G	AAAT	[AAG	ATA	AATTA	ATCA	TTT	AAAGO	GAG '	TGGT	AA		1076
			TTC Phe												GGC Gly		1124
			TCG Ser 20														1172
Glu	Thr	Met	GTG Val	Val	Thr	Ala	Ala	Gly	Tyr	Ala	Gln	Val					1220
			AGT Ser														1268
			GAT Asp														1316
			GGG Gly														1364
TCA Ser	AAC Asn	TAT Tyr	ACT Thr	CTT Leu	ATC Ile	TTG Leu	Val	GAT	GGT Gly	AAG Lyb	CGC Arg	Gln	ACC	TCA Ser	CGC Arg		1412

CAG ACC CGT CCA AAC AGC GAT GGC CCG GG Gln Thr Arg Pro Asn Ser Asp Gly Pro Gl 115		1460			
CCG CCA CTG CAA GCG ATT GAA CGT ATC GA Pro Pro Leu Gln Ala Ile Glu Arg Ile Gl 130	G GTG ATC CGT GGC CCG ATG u Val Ile Arg Gly Pro Met 140	1508			
TCT ACG CTG TAC GGC TCG GAT GCT GAC Ser Thr Leu Tyr Gly Ser Asp Ala Asp 145		1535			
(2) INFORMATION FOR SEQUENCE IDENTIFIC	ATION NUMBER: 2:				
(i) SEQUENCE CHARACTERISTICS:					
(A) LENGTH: (B) TYPE: (C) STRANDEDNESS: (D) TOPOLOGY:	51 nucleic acid both linear				
(xi) SEQUENCE DESCRIPTION: SEQ I	D NO: 2:				
TCCATGTGTC GATCTATCTC CAGTACAGAA TATATC	GAATA ATCCGCTTCT G	51			
(2) INFORMATION FOR SEQUENCE IDENTIFICA	ATION NUMBER: 3:				
(i) SEQUENCE CHARACTERISTICS:					
(A) LENGTH: (B) TYPE: (C) STRANDEDNESS: (D) TOPOLOGY:	120 nucleic acid both linear				
(xi) SEQUENCE DESCRIPTION: SEQ I	D NO: 3:				
AGATCTTGCA TAGGTATTTG ACCCTTAAAG AATAA	TTACC ACAGACGTTC CATATTTGGA	60			
CCGAACTATT CCATGTGTCG ATCTATCTCC AGTAC	AGAAT ATATGAATAA TCCGCTTCTG	120			
(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 4:					
(i) SEQUENCE CHARACTERISTICS:					
(A) LENGTH:(B) TYPE:(C) STRANDEDNESS:(D) TOPOLOGY:	158 nucleic acid both linear				
(xi) SEQUENCE DESCRIPTION: SEQ II	D NO: 4:				
AGATCTTGCA TAGGTATTTG ACCCTTAAAG AATAA	TTACC ACAGACGTTC CATATTTGGA	60			
CCGAACTATT CCATGTGTCG ATCTATCTCC AGTAC	AGAAT ATATGAATAA TCCGCTTCTG	120			
AAATTAAGAA TAATTATCAT TTAAAGGAGT GGTAAA					

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:	5:
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTE: 80 (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: linear	
' (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:	
CGATCTATCT CCAGTACAGA ATATATGAAT AATCCGCTTC TGAAATTAAG	AATAATTATC 60
ATTTAAAGGA GTGGTAAATG	80
(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:	6:
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 137 (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:	
AAGCTTTTAC GGCGCTGAGA TCTTGCATAG GTATTTGACC CTTAAAGAAT	
GACGITCCAT ATTTGGACCG AACTATTCCA TGTGTCGATC TATCTCCAGI	ACAGAATATA 120
TGAATAATCC GCTTCTG	137
(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:	7:
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 175 (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: linear	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:	
AAGCTTTTAC GGCGCTGAGA TCTTGCATAG GTATTTGACC CTTAAAGAAT	AATTACCACA 60
GACGTTCCAT ATTTGGACCG AACTATTCCA TGTGTCGATC TATCTCCAGT	ACAGAATATA 120
TGAATAATCC GCTTCTGAAA TTAAGAATAA TTATCATTTA AAGGAGTGGT	AAATG 175
(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER:	8:
(i) SEQUENCE CHARACTERISTICS:	
(A) LENGTH: 534 (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: linear	

(xi)	SEQUENCE	DESCRIPTION:	SEO	ID	NO:	8:

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GACGTTCCAT	ATTTGGACCG	AACTATTCCA	TGTGTCGATC	TATCTCCAGT	ACAGAATATA	120
TGAATAATCC	GCTTCTGAAA	TTAAGAATAA	TTATCATTTA	AAGGAGTGGT	AAATGTCCAG	180
ATTCAATCCA	TCCCCCGTCA	GTTTATCTGT	GACACTAGGC	TTAATGTTTT	CGGCTAGCGC	240
TTTTGCTCAA	GACGCGACGA	AAACGGATGA	AACCATGGTG	GTCACTGCGG	CGGGATACGC	300
GCAAGTGATT	CAAAATGCAC	CAGCCAGTAT	CAGTGTGATT	TCAAGAGAAG	ATCTGGAATC	360
TCGCTATTAC	CGTGATGTGA	CCGATGCGCT	AAAAAGCGTA	CCGGGTGTGA	CAGTCACCGG	420
AGGGGGCGAT	ACTACCGATA	TCAGCATTCG	TGGTATGGGA	TCAAACTATA	CTCTTATCTT	480
GGTGGATGGT	AAGCGCCAAA	CCTCACGCCA	GACCCGTCCA	AACAGCGATG	GCCC	534

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 9:

(i) SEQUENCE CHARACTERISTICS:

(A)	LENGTH:	517

(B) TYPE: nucleic acid

(C) STRANDEDNESS: both (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

AGATCTTGCA	TAGGTATTTG	ACCCTTAAAG	AATAATTACC	ACAGACGTTC	CATATTTGGA	60
CCGAACTATT	CCATGTGTCG	ATCTATCTCC	AGTACAGAAT	ATATGAATAA	TCCGCTTCTG	120
AAATTAAGAA	TAATTATCAT	TTAAAGGAGT	GGTAAATGTC	CAGATTCAAT	CCATCCCCCG	180
TCAGTTTATC	TGTGACACTA	GGCTTAATGT	TTTCGGCTAG	CGCTTTTGCT	CAAGACGCGA	240
CGAAAACGGA	TGAAACCATG	GTGGTCACTG	CGGCGGGATA	CGCGCAAGTG	ATTCAAAATG	300
CACCAGCCAG	TATCAGTGTG	ATTTCAAGAG	AAGATCTGGA	ATCTCGCTAT	TACCGTGATG	360
TGACCGATGC	GCTAAAAAGC	GTACCGGGTG	TGACAGTCAC	CGGAGGGGC	GATACTACCG	420
ATATCAGCAT	TCGTGGTATG	GGATCAAACT	ATACTCTTAT	CTTGGTGGAT	GGTAAGCGCC	480
AAACCTCACG	CCAGACCCGT	CCAAACAGCG	ATGGCCC			517

(2) INFORMATION FOR SEQUENCE IDENTIFICATION NUMBER: 10:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH:

nucleic acid

(B) TYPE: (C) STRANDEDNESS: (D) TOPOLOGY: both linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

WO 94/19482

- 31 -

CGATCTATCT	CCAGTACAGA ATATATO	AAT AATCCGCTTC TO	GAAATTAAG	AATAATTATC	60
ATTTAAAGGA	GTGGTAAATG TCCAGAT	TCA ATCCATCCCC CO	GTCAGTTTA	TCTGTGACAC	120
TAGGCTTAAT	GTTTTCGGCT AGCGCTT	TTG CTCAAGACGC G	ACGAAAACG	GATGAAACCA	180
TGGTGGTCAC	TGCGGCGGGA TACGCGC	AAG TGATTCAAAA TO	GCACCAGCC	AGTATCAGTG	240
TGATTTCAAG	AGAAGATCTG GAATCTC	GCT ATTACCGTGA TO	GTGACCGAT	GCGCTAAAAA	300
GCGTACCGGG	TGTGACAGTC ACCGGAG	GGG GCGATACTAC CO	GATATCAGC	ATTCGTGGTA	360
TGGGATCAAA	CTATACTCTT ATCTTGG	TGG ATGGTAAGCG CO	CAAACCTCA	CGCCAGACCC	420
GTCCAAACAG	CGATGGCCC				439
•	ATION FOR SEQUENCE SEQUENCE CHARACTERI (A) LENGTH:		JMBER:	11:	
(v i)	(B) TYPE: (C) STRANDEDNESS: (D) TOPOLOGY: SEQUENCE DESCRIPTI	nucl sind line	ar		
CCGAATTCTC	TAGAGATATC GTGTGGA	ATT GTGAGCGGAT AA	A	12:	42
(i) :	SEQUENCE CHARACTERI	STICS:			
	(A) LENGTH:(B) TYPE:(C) STRANDEDNESS:(D) TOPOLOGY:				
(xi)	SEQUENCE DESCRIPTION	ON: SEQ ID NO: 12	2:		
CCAAGCTTCT	GCAGCCCGGG ATTTAAC	מיים איים אויים איים איים איים איים איים	COT		A.E.

CLAIMS

- 1. A bacterial chromosome containing a DNA sequence encoding a heterologous antigen, said DNA sequence being functionally linked to an iron-regulated promoter.
- 5 2. The chromosome of claim 1, wherein said chromosome is a *Vibrio cholerae* chromosome.
- 3. The chromosome of claim 1, wherein said chromosome is a chromosome of an E. coli bacterium, a Shigella bacterium, a Salmonella bacterium, a Yersenia bacterium, a Citrobacter bacterium, an Enterobacter bacterium, a Klebsiella bacterium, a Proteus bacterium, a Providencia bacterium, a Serratia bacterium, a Vibrio bacterium, a Plesiomonas bacterium, an Aeromonas bacterium, or a bacille Calmette-Guerin (BCG).
- 15 4. The chromosome of claim 1, wherein said promoter is the promoter of a naturally-occurring V. cholerae gene.
- The chromosome of claim 4, wherein said promoter is the V. cholerae irgA promoter, and said
 chromosome lacks part or all of the irgA coding sequence.
 - 6. The chromosome of claim 5, wherein said promoter comprises a nucleotide sequence substantially identical to SEQ ID NO: 2.
- 7. The chromosome of claim 1, wherein said 25 heterologous antigen is a nontoxic polypeptide which induces an antigenic response in an animal.

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8. The chromosome of claim 7, wherein said polypeptide is a portion or all of a protein naturally expressed by an infectious organism.

- 9. The chromosome of claim 8, wherein said 5 infectious organism is a bacterium.
 - 10. The chromosome of claim 9, wherein said polypeptide is an immunogenic, nontoxic subunit or fragment of a bacterial toxin.
- 11. The chromosome of claim 10, wherein said
 10 toxin is Shiga toxin, diphtheria toxin, Pseudomonas
 exotoxin A, cholera toxin, pertussis toxin, tetanus
 toxin, anthrax toxin, E. coli heat-labile toxin (LT), E.
 coli heat-stable toxin (ST), or E. coli Shiga-like toxin.
- 12. The chromosome of claim 9, wherein said 15 protein is an OSP (Outer Surface Protein) of Borrelia burgdorferai.
 - 13. The chromosome of claim 8, wherein said infectious organism is a virus and said polypeptide is an immunogenic portion of a viral capsid.
- 20 14. The chromosome of claim 13, wherein said virus is a human immunodeficiency virus (HIV), one of the Herpes viruses, an influenza virus, a poliomyelitis virus, a measles virus, a mumps virus, or a rubella virus.
- 25 15. The chromosome of claim 8, wherein said infectious organism is a eukaryotic parasite.

- 34 -

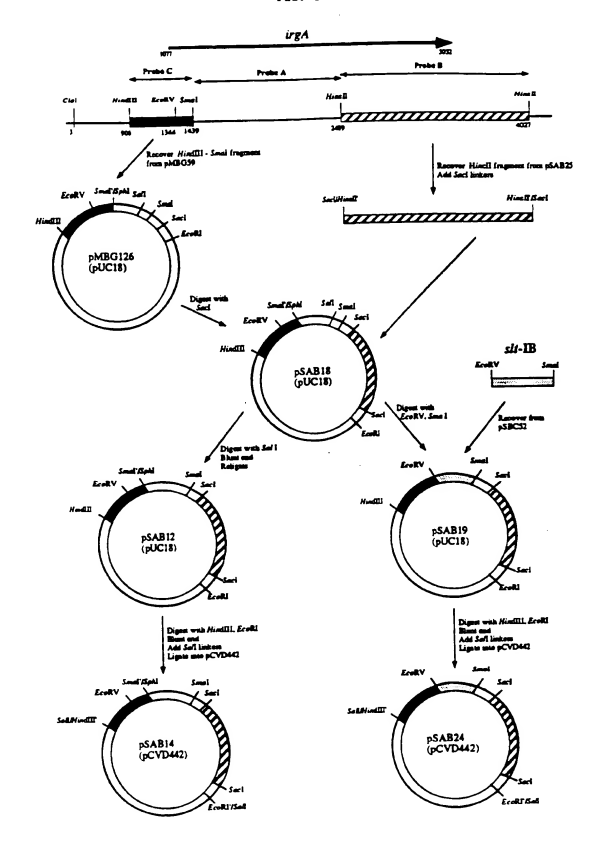
- 16. The chromosome of claim 15, wherein said parasite is the causative agent for malaria, pneumocystis pneumonia, or toxoplasmosis.
- 17. The chromosome of claim 16, wherein said 5 protein is a malarial circumsporozoite protein.
 - 18. The chromosome of claim 2, wherein said chromosome does not encode biologically active cholera toxin A subunit.
- 19. The chromosome of claim 5, wherein said10 chromosome does not encode biologically active cholera toxin A subunit.
- 20. A V. cholerae chromosome containing a DNA sequence encoding a heterologous antigen, said DNA sequence being functionally linked to a naturally15 occurring V. cholerae promoter.
- 21. The chromosome of claim 20, wherein said promoter is the promoter of a naturally-occurring gene encoding a V. cholerae virulence factor that is nonessential for growth of said cell, the coding sequence encoding said virulence factor being mutated or deleted so that said chromosome cannot express a biologically active form of said virulence factor.
 - 22. The chromosome of claim 20, wherein said promoter is the *irgA* promoter.
- 25 23. The chromosome of claim 20, wherein said heterologous antigen is part or all of a nontoxic polypeptide which is naturally expressed by an infectious

organism, which antigen induces an antigenic response in an animal.

- 24. The chromosome of claim 23, wherein said infectious organism is a bacterium.
- 5 25. The chromosome of claim 24, wherein said antigen is an immunogenic, nontoxic subunit or fragment of a bacterial toxin.
- 26. The chromosome of claim 25, wherein said toxin is Shiga toxin, diphtheria toxin, Pseudomonas
 10 exotoxin A, cholera toxin, pertussis toxin, tetanus toxin, anthrax toxin, E. coli LT, E. coli ST, or E. coli Shiga-like toxin.
- 27. The chromosome of claim 23, wherein said infectious organism is a virus and said antigen is an 15 immunogenic portion of a viral capsid.
- 28. The chromosome of claim 27, wherein said virus is a human immunodeficiency virus (HIV), one of the Herpes viruses, an influenza virus, a poliomyelitis virus, a measles virus, a mumps virus, or a rubella 20 virus.
 - 29. The chromosome of claim 23, wherein said infectious organism is a eukaryotic parasite.
- 30. The chromosome of claim 29, wherein said parasite is the causative agent for malaria, pneumocystis pneumonia, or toxoplasmosis.

- 36 -

- 31. The chromosome of claim 20, wherein said chromosome does not encode biologically active cholera toxin A subunit.
- 32. A V. cholerae cell, the chromosome of which 5 is the chromosome of claim 1.
 - 33. A $V.\ cholerae$ strain, the chromosome of which is the chromosome of claim 1.
 - 34. A homogeneous population of *V. cholerae* cells, each of which comprises the chromosome of claim 1.
- 35. A live-cell vaccine comprising the cell of claim 32 in a pharmaceutically acceptable diluent suitable for oral administration.
- 36. The vaccine of claim 35, wherein said chromosome does not encode biologically active cholera toxin A subunit.
 - 37. The vaccine of claim 36, wherein said chromosome does not encode biologically active IrgA.
 - 38. The vaccine of claim 37, wherein said heterologous antigen is Shiga-like toxin B subunit.
- 20 39. A method of vaccinating an animal comprising orally administering to said animal the vaccine of claim 35.
 - 40. The method of claim 39, wherein said animal is a human.



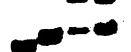
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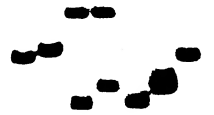
Fig. 2A 1 2 3 4 5 6 7 Fig. 2B 1 2 3 4 5 6 7

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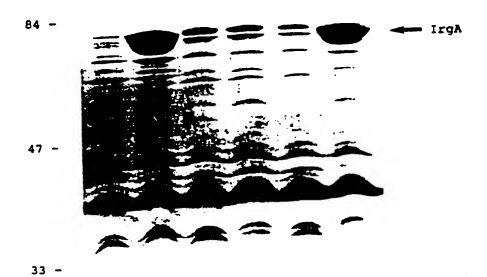
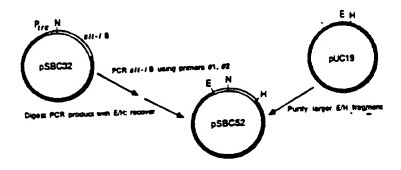


FIG. 4



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ational application No. INTERNATIONAL SEARCH REPORT PC 1/US94/01780 CLASSIFICATION OF SUBJECT MATTER IPC(5) :C12P-22-46; C12N 1720, 1/12; A01N 63/00 US CL :435/69.1, 252.3, 252.33, 253.1; 424/93A, 93P, 93D According to International Patent Classification (IPC) or to both national classification and IPC FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) U.S.: 435/69.1, 252.3, 252.33, 253.1; 424/93A, 93P, 93D Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) APS, DIALOG C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. Category* Υ WO, A, 89/02924 (BREY ET AL.) 06 APRIL 1989, see entire document. Y US, A, 4,882,278 (MEKALANOS) 21 NOVEMBER 1989, see 1-40 entire document. Y PNAS, USA, Volume 88, Number 4, issued 15 February 1-40 1991, Goldberg et al., "Positive transcriptional regulation of an iron-regulated virulence gene in Vibrio cholerae", pages 1125-1129, see entire document.

	Further documents are usted in the continuation of Box C	ــا	See patent family annex.
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C (Continu	tion) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	INFECTION AND IMMUNITY, Volume 58, No.1, issued January 1990, Goldberg et. al., "Identification of an Iron-regulated virulence determinant in Vibrio cholerae, using TnphoA mutants, pages 55-60, see entire document.	1-40
Y	NATURE, Volume 327, issued 11 JUNE 1987, Jacobs et al., "Introduction of foreign DNA into mycobacteria using a shuttle phasmid", pages 532-534, see entire document.	1-40